RNAscope® VS Universal AP Assay
For Ventana DISCOVERY™ ULTRA System

RED

Document Number 323250-USM-ULT
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Citing RNAscope® Assay in Publications

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Chapter 1. Product Information

About this guide

This user manual provides two versions of the RNAscope® VS Universal AP Assay:

- Appendix A. Semi-automated RNAscope® VS Universal AP Assay starting on page 23.

Product description

Background

The RNAscope® VS Universal Assays use a novel and proprietary method of in situ hybridization (ISH) to visualize single RNA molecules per cell in formalin-fixed, paraffin-embedded (FFPE) tissue mounted on slides. The assays are based on Advanced Cell Diagnostic’s patented signal amplification and background suppression technology, and can detect RNA molecules in archival samples and partially degraded specimens. The RNAscope® VS Universal Assay allows users to automate the highly sensitive RNAscope® Assay using the Ventana DISCOVERY™ ULTRA System.

Overview

Figure 1 on page 6 illustrates the RNAscope® VS Universal Assay procedure, which can be completed on the instrument in ~11 hours. Starting with properly prepared samples, sections are first pretreated, and then RNA-specific probes are hybridized to target RNA. The signal is amplified using multiple steps, followed by hybridization to horseradish peroxidase (HRP)- or alkaline phosphatase (AP)-labeled probes and detection using a chromogenic substrate. A single RNA transcript appears as a distinct dot of chromogen precipitate visible using a common bright-field microscope.
**Figure 1. Procedure overview**

1: Tissue section  
Start with properly prepared sections and pretreat to allow access to target RNA.

2: Hybridize to target RNA  
Hybridize gene-specific probe pairs to the target mRNA.

3: Amplify signal  
Probes are hybridized to a cascade of signal amplification molecules, culminating in binding of HRP- or AP-labeled probes. The Universal Assay enhances signal further with additional amplification steps. Add DAB or Fast Red substrate to detect target RNA.

4: Image  
Visualize target RNA using a standard bright field microscope.

**Kit contents and storage**

The RNAscope® VS Universal Assay requires the RNAscope® 2.5 VS Probes and the RNAscope® VS Universal Reagents, available from Advanced Cell Diagnostics.

**RNAscope® VS Probes**

The RNAscope® VS Probes consist of the user-specified Target Probe and the Positive and Negative Control Probes. Visit [https://acdbio.com/products](https://acdbio.com/products) to find a gene-specific Target Probe or appropriate Control Probes. Each probe is sufficient for staining ~30 standard slides. The probes have a shelf life of two years from the manufacturing date when stored as indicated in the following table:

<table>
<thead>
<tr>
<th>Target Probes</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Reagent</strong></td>
<td><strong>Cat. No.</strong></td>
</tr>
<tr>
<td>RNAscope® 2.5 VS Target Probe (species) – [gene]</td>
<td>Various</td>
</tr>
</tbody>
</table>

**Control Probes**

<table>
<thead>
<tr>
<th>Control Probes</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Reagent</strong></td>
<td><strong>Cat. No.</strong></td>
</tr>
<tr>
<td>RNAscope® 2.5 VS Positive Control Probe – [species] – PPIB</td>
<td>Various</td>
</tr>
<tr>
<td>RNAscope® 2.5 VS—— Negative Control Probe – DapB</td>
<td>312039</td>
</tr>
</tbody>
</table>

**RNAscope® VS Control Slides**

The RNAscope® VS Control Slides (Cat. No. 310045 for Human control slide, HeLa; Cat. No. 310023 for Mouse control slide, 3T3) contain FFPE cell pellets sectioned and mounted on slides. The control slides can be used for assay control with the RNAscope® 2.5 VS Positive Control Probe and RNAscope® 2.5 VS Negative Control Probe. The slides have a shelf life of nine months from the manufacturing date when stored at 2–8°C with desiccants.
**RNAscope® VS Universal Reagents**

RNAscope® VS Universal kits provide enough reagents to stain ~60 standard slides. You will receive two kits when you order the RNAscope® VS Universal AP Reagent Kit (Cat. No. 323250).

RNAscope® VS Universal Reagents include:

- RNAscope® VS Universal AP Detection Reagents (Cat. No. 323260)
- RNAscope® VS Universal Sample Prep Reagents (Cat. No. 323220)
- RNAscope® VS Accessory Kit (Cat. No. 320630)

The reagents are Ready-To-Use (RTU) and have a shelf life of nine months from the manufacturing date when stored as indicated in the following table:

<table>
<thead>
<tr>
<th>Cat. No.</th>
<th>Reagent</th>
<th>Quantity</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>322211</td>
<td>RNAscope® VS Universal AP AMP 1</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322212</td>
<td>RNAscope® VS Universal AP AMP 2</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322213</td>
<td>RNAscope® VS Universal AP AMP 3</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322214</td>
<td>RNAscope® VS Universal AP AMP 4</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322261</td>
<td>RNAscope® VS Universal AP AMP 5</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322262</td>
<td>RNAscope® VS Universal AP AMP 6</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322217</td>
<td>RNAscope® VS Universal AP AMP 7</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>322118</td>
<td>RNAscope® VS Protease</td>
<td>14 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cat. No.</th>
<th>Reagent</th>
<th>Quantity</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>323741</td>
<td>RNAscope® VS Universal Target Retrieval v2</td>
<td>10 mL x 2 bottles</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>323742</td>
<td>RNAscope® VS Universal Dewax</td>
<td>14 mL x 1 bottle</td>
<td>Room Temp (15–30°C)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cat. No.</th>
<th>Reagent</th>
<th>Quantity</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>320631</td>
<td>RNAscope® VS Hematoxylin — RTU</td>
<td>7 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
<tr>
<td>320632</td>
<td>RNAscope® VS Bluing Reagent — RTU</td>
<td>7 mL x 1 bottle</td>
<td>2–8°C</td>
</tr>
</tbody>
</table>

**IMPORTANT!**  Dewax must be in solution and at room temperature before use on the instrument. Warm in the hand, or place at 37°C for 15 MIN before each use regardless of the prior storage condition, since it may precipitate during shipment.

**IMPORTANT!**  Use only RNAscope® 2.5 VS Probes. Do not substitute the reagent components of the RNAscope® VS Universal Reagent Kit with those of other RNAscope® Reagent Kits.
Required materials from Roche Diagnostics

The RNAscope® VS Universal Assay requires specific materials and equipment available only from Roche Diagnostics (Ventana Medical Systems, Inc.). Catalog Numbers are valid in the U.S. only. For other regions, please check Catalog or ordering numbers with your local lab supplier.

### Probe Dispensers (Cat. No. 960-761 to 960-780; for Ordering Code, please contact local Roche representative)

<table>
<thead>
<tr>
<th>Component</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>250 Test Probe #1–20 dispensers — fill dispensers with RNAscope® 2.5 VS Probes. Use up to 20 probes at one time.</td>
<td>Room Temp (15–30°C)</td>
</tr>
</tbody>
</table>

### mRNA Sample Prep Kit (Cat. No. 760-248; Ordering Code 08127166001)

<table>
<thead>
<tr>
<th>Component</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>mRNA Target Retrieval dispenser — fill dispenser with RNAscope® VS Universal Target Retrieval v2</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA Dewax — fill dispenser with RNAscope® VS Universal Dewax</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA Protease dispenser — fill dispenser with RNAscope® VS Protease</td>
<td>Room Temp (15–30°C)</td>
</tr>
</tbody>
</table>

### mRNA RED Probe Amplification Kit (Cat. No. 760-236; Ordering Code 7095341001)

<table>
<thead>
<tr>
<th>Component</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>mRNA AMP 1 dispenser — fill dispenser with VS Universal AP AMP 1</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA AMP 2 dispenser — fill dispenser with VS Universal AP AMP 2</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA AMP 3 dispenser — fill dispenser with VS Universal AP AMP 3</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA AMP 4 dispenser — fill dispenser with VS Universal AP AMP 4</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA AMP 5 dispenser — fill dispenser with VS Universal AP AMP 5</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA AMP 6 dispenser — fill dispenser with VS Universal AP AMP 6</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>mRNA AMP 7 dispenser — fill dispenser with VS Universal AP AMP 7</td>
<td>Room Temp (15–30°C)</td>
</tr>
</tbody>
</table>

### mRNA RED Detection Kit (Cat. No. 760-234; Ordering Code 7099037001)

<table>
<thead>
<tr>
<th>Component</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>mRNA Inhibitor-prefilled</td>
<td>2–8°C</td>
</tr>
<tr>
<td>mRNA Activator dispenser-prefilled</td>
<td>2–8°C</td>
</tr>
<tr>
<td>mRNA Napthol dispenser-prefilled</td>
<td>2–8°C</td>
</tr>
<tr>
<td>mRNA Fast Red dispenser-prefilled</td>
<td>2–8°C</td>
</tr>
</tbody>
</table>

### Generic Dispensers (Cat. No. 771-741; Ordering Code 05271720001, Cat. No. 771-742; Ordering Code 05271738001)

<table>
<thead>
<tr>
<th>Component</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>250 Test Counterstain 1 dispenser — fill dispenser with VS Hematoxylin</td>
<td>Room Temp (15–30°C)</td>
</tr>
<tr>
<td>250 Test Counterstain 2 dispenser — fill dispenser with VS Bluing Reagent</td>
<td>Room Temp (15–30°C)</td>
</tr>
</tbody>
</table>
Equipment and buffers

<table>
<thead>
<tr>
<th>Component</th>
<th>Cat. No./Ordering Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X DISCOVERY Wash (RUO)</td>
<td>950-510 / 07311079001</td>
</tr>
<tr>
<td>ULTRA LCS (Predilute)</td>
<td>650-210 / 05424534001</td>
</tr>
<tr>
<td>SSC (10X)</td>
<td>950-110 / 05353947001</td>
</tr>
<tr>
<td>Reaction Buffer (10X)</td>
<td>950-300 / 05353955001</td>
</tr>
<tr>
<td>DISCOVERY CC1</td>
<td>950-500 / 06414575001</td>
</tr>
</tbody>
</table>

**IMPORTANT!** To run the VS Universal assay successfully, use DISCOVERY Wash (950-510). Do not use DISCOVERY EZ Prep. Place 2X SSC (950-110) in the SSC bulk container instead of Ribowash. You may fill the option bulk container with reaction buffer.

User-supplied materials

**IMPORTANT!** Do not substitute other materials for the SuperFrost® Plus Slides listed in the following table.

<table>
<thead>
<tr>
<th>Description</th>
<th>Supplier</th>
<th>Cat. No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>100% ethanol (EtOH)</td>
<td>American Master Tech Scientific/MLS*</td>
<td>ALREAGAL</td>
</tr>
<tr>
<td>xylene</td>
<td>Fisher Scientific/MLS</td>
<td>X3P-1GAL</td>
</tr>
<tr>
<td>10% neutral-buffered formalin (NBF)</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Paraffin wax</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>1X PBS</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Microtome</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Drying oven, capable of holding temperature at 60 +/- 1°C</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>EcoMount</td>
<td>Biocare</td>
<td>EM897L</td>
</tr>
<tr>
<td>Tissue-Tek® Vertical 24 Slide Rack</td>
<td>American Master Tech Scientific/MLS</td>
<td>LWSRA24</td>
</tr>
<tr>
<td>Tissue-Tek® Staining Dishes</td>
<td>American Master Tech Scientific/MLS</td>
<td>LWT4457EA</td>
</tr>
<tr>
<td>Tissue-Tek® Clearing Agent Dishes, xylene resistant</td>
<td>American Master Tech Scientific/MLS</td>
<td>LWT4456EA</td>
</tr>
<tr>
<td>Cover Glass 24 x 50 mm</td>
<td>Fisher Scientific/MLS</td>
<td>12-545F</td>
</tr>
<tr>
<td>Distilled water</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Dawn detergent or similar detergent</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Fume hood</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Optional: Glass beaker (1 or 2 L)</td>
<td>MLS</td>
<td>—</td>
</tr>
<tr>
<td>Optional: Hot plate</td>
<td>Fisher Scientific/MLS</td>
<td>11-300-49SHP</td>
</tr>
</tbody>
</table>

* Major Laboratory Supplier in North America. For other regions, please check Catalog Numbers with your local lab supplier.
Chapter 2. Before You Begin

Prior to running the RNAscope® VS Universal Assay on your samples for the first time, we recommend that you:

- Be familiar with the Ventana™ DISCOVERY™ ULTRA system. Refer to the Ventana™ System User Manual.
- Run the assay on FFPE RNAscope® VS Control Slides (Cat. No. 310045 for Human control slide, Hela; Catalog No. 310023 for Mouse control slide, 3T3) using the Positive and Negative Control Probes.

**Important procedural guidelines**

- Start with properly fixed and prepared sections. Refer to Chapter 3. Prepare and Pretreat Samples on page 11, Recommended guidelines on page 20, and to our sample preparation and pretreatment user guides available at https://acdbio.com/technical-support/user-manuals.
- Regularly maintain and clean your automated staining instrument.
- Always run positive and negative control probes on your sample to assess sample RNA quality and optimal permeabilization.
- Do not substitute required materials. Assay has been validated with these materials only.
- Follow the protocol exactly for best results.
- Do not let your sections dry out during the procedure unless specified in the protocol.
- Use good laboratory practices and follow all necessary safety procedures. Refer to Appendix B. Safety on page 32 in this document for more information.
Chapter 3. Prepare and Pretreat Samples

Formalin-fixed, paraffin-embedded (FFPE) sample preparation and pretreatment are described in the following protocols.

**IMPORTANT!** We highly recommend following these guidelines. We cannot guarantee assay results with other preparation methods.

For samples treated differently from the following protocol, you may need to optimize pretreatment conditions. Refer to Recommended guidelines on page 20, and to https://acdbio.com/technical-support/solutions.

### Prepare FFPE sections

**Materials required**

- 10% neutral buffered formalin (NBF)
- 1X PBS
- Paraffin wax
- 100% ethanol (EtOH)
- xylene
- Microtome
- Water bath
- SuperFrost® Plus slides

**Fix the sample**

1. Immediately following dissection, fix tissue in 10% NBF for **16–32 HRS** at ROOM TEMPERATURE (RT).
   
   Fixation time will vary depending on tissue type and size.

   **CAUTION!** Handle biological specimens appropriately.

   **IMPORTANT!** Fixation for **<16 HRS** or **>32 HRS** will impair the performance of the RNAscope® VS Universal Assay.

**Dehydrate, embed, and cut the sample**

**IMPORTANT!** Use fresh reagents.

1. Wash sample with 1X PBS.
2. Dehydrate sample using a standard ethanol series, followed by xylene.
3. Embed sample in paraffin using standard procedures.

   **Note:** Embedded samples may be stored at 15–25°C with desiccation. To better preserve RNA quality over a long period (>1 yr), storing at 2–8°C with desiccation is recommended.

4. Trim paraffin blocks as needed, and cut embedded tissue into 5 +/– 1 μm sections using a microtome.
5. Place paraffin ribbon in a 40–45°C water bath, and mount sections on SUPERFROST® PLUS SLIDES. Place tissue as shown for optimal staining:

![Diagram showing tissue section location]

**IMPORTANT!** Do not mount more than one section per slide. Place sections in the center of the slide.

6. Air dry slides **OVERNIGHT** at RT. Do NOT bake slides unless they will be used for RNAscope® within 1 week.

**OPTIONAL STOPPING POINT.** Use sectioned tissue within 3 months. Store sections with dessicants at RT.
Chapter 4. Automated RNAscope® VS Universal AP Assay

**IMPORTANT!** We strongly recommend you run the RNAscope® VS Control Slides (Cat. No. 310045 or 310023) using the RNAscope® 2.5 VS Positive and Negative Control Probes along with your samples in every run.

Appendix A. Semi-automated RNAscope® VS Universal AP Assay on page 23 describes an offline boiling procedure for use with Cat. No.322000.

**Workflow**

1. Prepare the materials
2. Run the RNAscope® VS Universal AP Assay ~ 11 HRS
3. Wash and mount slides
Prepare the materials

Materials required

<table>
<thead>
<tr>
<th>Materials Provided by Advanced Cell Diagnostics</th>
<th>Materials Provided by Ventana™ Medical Systems</th>
<th>Other Materials and Equipment</th>
</tr>
</thead>
<tbody>
<tr>
<td>• RNAscope® 2.5 VS Target Probe</td>
<td>• DISCOVERY™ ULTRA — automated slide stainer</td>
<td>• Distilled water</td>
</tr>
<tr>
<td>• RNAscope® 2.5 VS Positive Control Probe</td>
<td>• DISCOVERY Wash Buffer 10X</td>
<td>• Dawn detergent or similar</td>
</tr>
<tr>
<td>• RNAscope® 2.5 VS Negative Control Probe</td>
<td>• ULTRA LCS (Predilute)</td>
<td>detergent</td>
</tr>
<tr>
<td>• RNAscope® VS Universal Dewax</td>
<td>• SSC Buffer 10X</td>
<td>• Fume hood</td>
</tr>
<tr>
<td>• RNAscope® VS Protease</td>
<td>• DISCOVERY CC1</td>
<td>• xylene</td>
</tr>
<tr>
<td>• RNAscope® VS Target Retrieval v2</td>
<td>• Reaction Buffer 10X</td>
<td>• Tissue-Tek® Staining Dish</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 1</td>
<td>• Probe dispensers</td>
<td>• Tissue-Tek® Clearing Agent</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 2</td>
<td>• mRNA RED Probe Amplification Kit</td>
<td>• Dish, xylene-resistant</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 3</td>
<td>• mRNA Universal Sample Prep Kit</td>
<td>• Tissue-Tek® Vertical 24 Slide</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 4</td>
<td>• mRNA RED Detection Kit</td>
<td>• Rack</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 5</td>
<td>• User fillable dispensers</td>
<td>• EcoMount</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 6</td>
<td></td>
<td>• Cover Glass, 24 mm x 50 mm</td>
</tr>
<tr>
<td>• RNAscope® VS Hematoxylin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• RNAscope® VS Bluing Reagent</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Prepare the instrument

If the instrument has not been used for ≥1 week, follow the guidelines for instrument maintenance in the Ventana™ System User Manual.

Dilute bulk reagents

Please prepare the bulk fluids as per manufacturer’s instructions.

Register new reagents

Reagent dispensers come with appropriate barcode labels and registration buttons for dispensing RNAscope® VS Universal AP Reagents. Refer to the Ventana™ DISCOVERY ULTRA System User Manual for details. To register reagents:

1. Log all ACD reagents and probes into the software as “log user-fillable reagents” or “log user-fillable probes”.
2. Use the wand that comes with the instrument to register new reagent kits.

Prepare instrument reagents

Refer to the table on page 8 to determine the proper dispenser for each reagent.

1. For RNAscope® VS Universal AP reagents AMP 1 to AMP 7, transfer the entire volume of each component into the correspondingly labeled dispenser.
2. Transfer the 2.5 VS Target Probe, 2.5 VS Positive Control Probe, 2.5 VS Negative Control Probe, VS Universal Dewax, VS Protease, both bottles of VS Target Retrieval v2, VS Hematoxylin, and VS Bluing Reagent to the correspondingly labeled dispenser (see page 8).
IMPORTANT! Avoid cross contamination between reagents. Dewax must be warmed to room temperature and be completely in solution before use.

3. Follow the dispenser product insert instructions to properly prime and handle the dispensers.
4. Store tightly-capped dispensers (except the Dewax dispenser) at 4°C when not in use.

IMPORTANT! Do not use expired reagents.

5. Empty the waste bottle if needed.

Create an instrument protocol

1. Open the VS software and click on the Protocol button.
2. Click on Create/Edit Protocols, go to the Procedure drop down menu and select mRNA Universal.
3. Main protocol steps appear as shown:

4. After selecting the main protocol steps, drop down menus become available. Select the appropriate protocol steps by clicking on the associated check boxes as shown:

Note: The length of time that the tissue undergoes Cell Conditioning is equal to the checked box with the longest time. It is not cumulative of all checked times.

Note: Make sure you select mRNA AP Detection.
5. Select the appropriate assay conditions from the drop down menus according to the following table:

<table>
<thead>
<tr>
<th>Tissue Type</th>
<th>Cell Conditioning Temperature</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain</td>
<td>97°C</td>
<td>16 MIN</td>
</tr>
<tr>
<td>Cell pellet</td>
<td>97°C</td>
<td>16 MIN</td>
</tr>
<tr>
<td>Colon</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Heart</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Intestine</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Kidney</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Liver</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Lung</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Prostate</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Spleen</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Tonsil</td>
<td>97°C</td>
<td>24 MIN</td>
</tr>
<tr>
<td>Brain</td>
<td>97°C</td>
<td>16 MIN</td>
</tr>
</tbody>
</table>

**Standard Temperatures/Times**

<table>
<thead>
<tr>
<th>VS Protease</th>
<th>37°C</th>
</tr>
</thead>
</table>
| Standard probe temperatures | Single Probes 43°C  
Pooled Probes 50°C |
| Standard AMP 1 and AMP 2 temperatures* | 39°C |
| AMP 5 incubation time** | 4 MIN |

* For very high backgrounds, you can raise temperatures to 40 or 41°C. However, some signal may be lost. Please contact support or your FAS before making these changes.

6. Click **Save as**, then select a protocol number from the drop down menu and choose a protocol name for each probe. Click **Save**.

7. Click **Close** to go back to the main screen.

8. Assign probe number from the list to each probe of interest. For each probe selected, assign a protocol.

**Print the labels**

1. Select the **Print Label** icon from the upper right corner of the home page screen.
2. Select your preferred template or create a new template. To create a new template, refer to the Ventana™ DISCOVERY ULTRA System User Manual for details.
3. Click on **Protocol**.
4. Select the protocol you created for the RNAscope® VS Universal AP Assay and click on the **Add** button. When the protocols for all of the slides have been assigned, click on **Close/Print**.
5. Fill in the template for each slide. Select **Print** when completed.
6. Proceed to the following procedure **Load the reagents**.
Run the RNAscope® VS Universal AP Assay

Materials required

- Prepared slides
- Prepared instrument reagents
- Prepared detergent
- Distilled water
- Prepared dehydrating materials
- Tissue-Tek® Vertical 24 Slide Rack
- Tissue-Tek® Staining Dish
- EcoMount
- Cover Glass, 24 mm x 50 mm
- Fume hood
- Xylene

Load the reagents

1. Remove the nozzle caps from the filled dispensers and place each cap on the post located on the back of the dispenser.
2. If needed, remove any air bubbles at the nozzle tips by pushing down on the nozzle until the liquid reaches the tip of the nozzle or forms a small meniscus at the tip of the nozzle.
   **IMPORTANT!** Do not dispense any drops as this could compromise your drop inventory.
3. Load dispensers onto the reagent racks.
4. Remove the yellow locking ring from the dispensers in the prefilled mRNA Red Detection Kit. Refer to the instructions provided by Ventana™ Medical Systems.
5. Load the reagent racks onto the reagent carousel.

Start the run

1. Click the **Ready** button.
2. Eject slide drawers.
3. Load each slide onto a heater pad with the label facing upward and inward. Ensure that the slides sit securely on the pads.
   **IMPORTANT!** Prior to loading the slides, ensure heater pads are completely dry. Wipe off any solution using laboratory tissue paper.
5. Click the Running button. Automated assay will finish in ~11 HRS.

**IMPORTANT!** Before leaving the instrument unattended, ensure all reagents and slides are successfully registered and the instrument is running.

### Prepare detergent

1. Prepare 200 mL of diluted detergent by adding 1-2 drops detergent to 200 mL distilled water in a container with a cap.
2. Mix well by inverting the container 4–5 times.
3. Add diluted detergent to a Tissue-Tek® Staining Dish.
   
   **Note:** Store diluted detergent at RT.

### Prepare dehydrating reagents

- In a fume hood, fill a clearing agent dish with ~200 mL fresh xylene.

**Note:** Ensure all containers remain covered when not in use.

### Complete the run

1. After the run is complete, remove the Dewax (Pretreatment A), place nozzle caps on the dispensers, and store at room temperature.
2. For the remaining reagents, place nozzle caps back on the dispensers and place racks onto magnet locking tray.

**IMPORTANT!** Store reagent racks at 4°C until next use. Store the Dewax dispenser at room temperature.

### Wash the slides

1. Submerge a Tissue-Tek® Slide Rack into the Tissue-Tek® Staining Dish containing 200 mL diluted detergent.
2. Open the instrument slide drawers and unload slides.
3. Decant solution on the slides into the slide drawer, then immediately load slides into the Tissue-Tek® Slide Rack submerged in detergent.
4. Rinse oil off the slides by moving the slide rack up and down in the dish 10 times.
5. Replace the detergent with distilled water and rinse slides by moving the slide rack up and down a minimum of 10 times.
6. Repeat Step 5 three to five times.

### Mount the samples

1. Remove the slide rack from the staining dish and dry slides in a 60°C dry oven for 30 MIN.
   
   **IMPORTANT!** The Red substrate is alcohol sensitive. Do not dehydrate the slides in alcohol.

2. Cool the slides for 5 MIN at RT.
3. Briefly dip one slide into into fresh pure xylene and immediately place 1–2 drops of EcoMount on the slide before the xylene dries.
IMPORTANT! Use the EcoMount mounting medium only.

4. Carefully place 24 mm x 50 mm coverslip over the tissue section. Avoid trapping air bubbles.
5. Repeat steps 2 and 3 for each slide.
6. Air dry slides for at least 5 MIN.
7. Proceed to Chapter 5. Evaluate the Results on page 21.

Recommended guidelines

We highly recommend following the guidelines for Cell Conditioning (Target Retrieval), and AP Detection 3rd Pretreatment (Protease) conditions for:

- Any new or previously untested FFPE tissue types.
- Samples prepared differently than the protocol in Chapter 3. Prepare and Pretreat Samples on page 11.

1. Stain six representative slides using the positive and negative control probes according to the following table:

<table>
<thead>
<tr>
<th>Slide No.</th>
<th>Probe</th>
<th>Target Retrieval</th>
<th>Protease</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Positive control</td>
<td>16 MIN</td>
<td>24 MIN</td>
</tr>
<tr>
<td>2</td>
<td>Negative control</td>
<td>16 MIN</td>
<td>24 MIN</td>
</tr>
<tr>
<td>3</td>
<td>Positive control</td>
<td>16 MIN</td>
<td>16 MIN</td>
</tr>
<tr>
<td>4</td>
<td>Negative control</td>
<td>16 MIN</td>
<td>16 MIN</td>
</tr>
<tr>
<td>5</td>
<td>Positive control</td>
<td>24 MIN</td>
<td>16 MIN</td>
</tr>
<tr>
<td>6</td>
<td>Negative control</td>
<td>24 MIN</td>
<td>16 MIN</td>
</tr>
</tbody>
</table>

2. Evaluate staining and tissue morphology as in Chapter 5. Evaluate the Results and determine which pretreatment condition yielded the highest positive control signal and lowest negative control signal. Using PPIB, positive control signal should have a staining score of 3 or higher, and the negative control signal should be 0.

3. Use the optimized pretreatment conditions to run the assay with the target probe.

4. If none of the conditions are satisfactory, contact technical support at support.acd@bio-techne.com.
Chapter 5. Evaluate the Results

Examine tissue sections under a standard bright field microscope at 20–40X magnification:

- Assess tissue and cell morphology.
- Assess positive control signal strength. Positive control signal should be visible as punctate dots within the cell cytoplasm at 20–40X magnification.
- Assess negative control background. One dot to every 10 cells displaying background staining per 40X microscope field is acceptable.
- Evaluate target probe signal using the scoring guidelines in the next section.

Scoring guidelines

The RNAscope® Assay enables a semi-quantitative scoring guideline utilizing the estimated number of punctate dots present within each cell boundary.

An example of how to develop such a guideline for semi-quantitative assessment of RNAscope® staining intensity is presented below for a gene with expression level varying between 1 to > 10 copies per cell.

Note: If your gene expression level is higher or lower than this range, you may need to scale the criteria accordingly.

Categorize staining into five grades: 0, 1+, 2+, 3+, and 4+ according to the following table:

<table>
<thead>
<tr>
<th>Staining Score</th>
<th>Microscope Objective Scoring*</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No staining, or less than 1 dot/10 cells (40X magnification)</td>
</tr>
<tr>
<td>1</td>
<td>1–3 dots/cell (visible at 20–40X magnification)</td>
</tr>
<tr>
<td>2</td>
<td>4–9 dots/cell. Very few dot clusters (visible at 20–40X magnification)</td>
</tr>
<tr>
<td>3</td>
<td>10-15 dots/cell and/or &lt;10% positive cells have dots in clusters (visible at 20X magnification)</td>
</tr>
<tr>
<td>4</td>
<td>&gt;15 dots/cell and/or &gt;10% positive cells have dots in clusters (visible at 20X magnification)</td>
</tr>
</tbody>
</table>

* Discount cells with artificially high nuclear background staining.

Quantitative image analysis

RNAscope® Spot Studio Software is designed for pathologists with no prior training in image analysis. This intuitive software allows users to get statistical results with complete information of cell-count/region and number of spots per cell. Simply load any image, select a region of interest, define settings and run analysis, followed by a quality control review before results are exported. Further information is available on our website at www.acdbio.com.
Troubleshooting

For troubleshooting information, please contact technical support at support.acd@bio-techne.com.

Tissue example

If the assay is successful, the staining should look like the following images:

Figure 2. RNAscope® VS Universal AP Assay results in HeLa cells

Hs-TBP (Positive Control)  DapB (Negative Control)
Appendix A. Semi-automated RNAscope® VS Universal AP Assay

Most sample types can be fully automated on the Discovery ULTRA. Manual pretreatment may give a better result in some cases. Use the semi-automated procedure for tissues that do not have a satisfactory result when using the fully automated procedure. See Chapter 4. Automated RNAscope® VS Universal AP Assay on page 13.

Workflow

<table>
<thead>
<tr>
<th>Prepare the materials</th>
</tr>
</thead>
<tbody>
<tr>
<td>↓</td>
</tr>
<tr>
<td>Bake the slides</td>
</tr>
<tr>
<td>↓</td>
</tr>
<tr>
<td>Deparaffinize FFPE sections</td>
</tr>
<tr>
<td>↓</td>
</tr>
<tr>
<td>Pretreat the slides</td>
</tr>
<tr>
<td>↓</td>
</tr>
<tr>
<td>Run the RNAscope® VS Universal AP Assay ~8 HRS</td>
</tr>
<tr>
<td>↓</td>
</tr>
<tr>
<td>Wash and mount slides</td>
</tr>
</tbody>
</table>

RNAscope® VS Universal AP Assay for the DISCOVERY® ULTRA System User Manual

323250-USM-ULT/Rev A/ Date: 06112019
Kit contents and storage

<table>
<thead>
<tr>
<th>Cat. No.</th>
<th>Reagent</th>
<th>Quantity</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>322000</td>
<td>RNAscope® Target Retrieval Reagents*</td>
<td>70 mL x 4 bottles</td>
<td>Room Temp (15–30°C)</td>
</tr>
</tbody>
</table>

* Not provided with the kit and needs to be purchased separately.

IMPORTANT! Do not substitute the reagent components of the RNAscope® VS Universal Reagent Kit with those of other RNAscope® Reagent Kits, even those having the same name. The Target Retrieval solution in the RNAscope® VS Universal Kit CANNOT be used for offline boiling. Please separately purchase the RNAscope® Target Retrieval Reagents (Cat. No. 322000) to boil samples off the instrument.

Prepare the materials

Materials can be prepared ahead of time or while baking the slides, unless otherwise stated. See Bake the slides on page 27.

Materials required

<table>
<thead>
<tr>
<th>Materials provided by Advanced Cell Diagnostics</th>
<th>Materials Provided by Ventana™ Medical Systems</th>
<th>Other Materials and Equipment</th>
</tr>
</thead>
<tbody>
<tr>
<td>• RNAscope® 2.5 VS Target Probe</td>
<td>• DISCOVERY™ ULTRA — automated slide stainer</td>
<td>• Distilled water</td>
</tr>
<tr>
<td>• RNAscope® 2.5 VS Positive Control Probe</td>
<td>• DISCOVERY Wash 10x</td>
<td>• Glass beaker (1 or 2 L)</td>
</tr>
<tr>
<td>• RNAscope® 2.5 VS Negative Control Probe</td>
<td>• ULTRA LCS (Predilute)</td>
<td>• Hot plate</td>
</tr>
<tr>
<td>• RNAscope® Target Retrieval Reagents (Offline)</td>
<td>• Reaction Buffer 10X</td>
<td>• Dawn detergent or similar detergent</td>
</tr>
<tr>
<td>• RNAscope® VS Protease</td>
<td>• Probe dispensers</td>
<td>• Fume hood</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 1</td>
<td>• mRNA RED Probe Amplification Kit</td>
<td>• xylene</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 2</td>
<td>• mRNA RED Detection Kit</td>
<td>• 100% ethanol (EtOH)</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 3</td>
<td>• User fillable dispensers</td>
<td>• Tissue-Tek® Staining Dishes</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 4</td>
<td>• mRNA Universal Sample Prep Kit</td>
<td>• Tissue-Tek® Clearing Agent Dishes, xylene-resistant</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 5</td>
<td>• CCI Buffer</td>
<td>• Tissue-Tek® Vertical 24 Slide Rack</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 6</td>
<td></td>
<td>• EcoMount</td>
</tr>
<tr>
<td>• RNAscope® VS Universal AP AMP 7</td>
<td></td>
<td>• Cover Glass, 24 mm x 50 mm</td>
</tr>
<tr>
<td>• RNAscope® VS Hematoxylin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• RNAscope® VS Bluing Reagent</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Prepare the instrument

If the instrument has not been used for ≥1 week, follow the guidelines for instrument maintenance in the Ventana™ System User Manual.

Dilute bulk reagents

Please prepare the bulk fluids as per manufacturer’s instructions.
Register new reagents

Reagent dispensers come with appropriate barcode labels and registration buttons for dispensing RNAscope® VS Reagents. Refer to the Ventana™ DISCOVERY ULTRA System User Manual for details.
To register reagents:

1. Log all ACD reagents and probes into the software as “log user-fillable reagents” or “log user-fillable probes”.
2. Use the wand that comes with the instrument to register new reagent kits.

Prepare instrument reagents

Refer to the table on page 8 to determine the proper dispenser for each reagent.

1. For RNAscope® VS Universal AP AMP 1 to AMP 7, transfer the entire volume of each component into the correspondingly labeled dispenser.
2. Transfer the 2.5 VS Target Probe, 2.5 VS Positive Control Probe, 2.5 VS Negative Control Probe, VS Protease, VS Hematoxylin, and VS Bluing Reagent to the correspondingly labeled dispensers.
3. Follow the dispenser product insert instructions to properly prime and handle the dispensers.
4. Store tightly-capped dispensers at 4°C when not in use.
5. Check solution levels: DISCOVERY Wash, 2X SSC, Reaction Buffer, ULTRA LCS (Predilute), and CC1. Refill if they are less than half full. To run a full tray of 30 slides, fully fill the 2X SSC and Reaction Buffer containers.

**IMPORTANT!** Do not use expired reagents.

6. Empty the waste carboy if needed.

Prepare deparaffinization reagents

- In a fume hood, fill two clearing agent dishes with ~200 mL fresh xylene.
- In a fume hood, fill two staining dishes with ~200 mL fresh 100% EtOH.

**Note:** Ensure all containers remain covered when not in use.

Prepare 1X Target Retrieval

Prepare 1X Target Retrieval while FFPE slides are baking at 60°C, or the following day if you choose the optional stopping point on page 27. 1X Target Retrieval is used in manual cell conditioning (CC).

1. Prepare 700 mL of fresh 1X Target Retrieval by adding 630 mL distilled water to 1 bottle (70 mL) 10X 1X Target Retrieval solution in the beaker.
2. Mix well and cover the beaker with foil.

**IMPORTANT!** Do not use RNAscope® VS Universal Target Retrieval v2 for offline boiling.

Create an instrument protocol

1. Open the VS software and click on the Protocol button.
2. Click on Create/Edit Protocols, go to the Procedure drop down menu and select mRNA Universal.
3. Main protocol steps appear as shown:

![Protocol Steps Diagram]

**IMPORTANT!** Do not select Baking, Deparaffinization, or Cell Conditioning.

4. After selecting the main protocol steps, drop down menus become available. Select the appropriate protocol steps by clicking on the associated check boxes as shown above.

5. Select the appropriate assay conditions from the drop down menus according to the following table:

<table>
<thead>
<tr>
<th></th>
<th><strong>Standard Temperatures/Times</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VS Protease</strong></td>
<td>37°C</td>
</tr>
<tr>
<td><strong>Standard probe temperatures</strong></td>
<td></td>
</tr>
<tr>
<td>Single Probes</td>
<td>43°C</td>
</tr>
<tr>
<td>Pooled Probes</td>
<td>50°C</td>
</tr>
<tr>
<td><strong>Standard AMP 1 and AMP 2 temperatures</strong>*</td>
<td>39°C</td>
</tr>
<tr>
<td><strong>AMP 5 incubation time</strong></td>
<td>4 MIN</td>
</tr>
</tbody>
</table>

* For very high backgrounds, you can raise temperatures to 40 or 41°C. However, some signal may be lost. Please contact support or your FAS before making these changes.

6. Click **Save As**, then select a protocol number from the drop down menu and choose a protocol name for each probe. Click **Save**.

7. Click **Close** to go back to the main screen.

8. Assign a probe number from the list to each probe of interest. For each probe selected, assign a protocol.

**Print the labels**

1. Select the **Print Label** icon from the bottom of the home page screen.
2. Select your preferred template or create a new template. To create a new template, refer to the **Ventana™ System User Manual** for details.
3. Select the protocol you created for the RNAscope® VS Universal Assay.
4. Click on **Protocol** to add and print the label.
Manually pretreat the samples

Materials required

<table>
<thead>
<tr>
<th>Materials Provided by the Target Retrieval Reagents (Cat. No. 322000)</th>
<th>Other Materials and Equipment</th>
</tr>
</thead>
<tbody>
<tr>
<td>• RNAscope® Target Retrieval Reagents</td>
<td>• Drying oven</td>
</tr>
<tr>
<td></td>
<td>• FFPE slides</td>
</tr>
<tr>
<td></td>
<td>• Tissue-Tek® Vertical 24 Slide Rack</td>
</tr>
<tr>
<td></td>
<td>• Tissue-Tek® Staining Dish</td>
</tr>
<tr>
<td></td>
<td>• Distilled water</td>
</tr>
<tr>
<td></td>
<td>• Prepared deparaffinization materials</td>
</tr>
<tr>
<td></td>
<td>• Glass beaker (1 or 2 L)</td>
</tr>
<tr>
<td></td>
<td>• Hot plate</td>
</tr>
</tbody>
</table>

Bake the slides

1. Bake slides in a dry oven for 30-60 MIN at 60°C.

OPTIONAL STOPPING POINT Use immediately or store at RT with desiccants for ≤1 week. Prolonged storage may degrade sample RNA.

IMPORTANT! If you continue, prepare the materials for the following protocols while the slides are baking: Deparaffinize FFPE sections, Pretreat the slides, and Run the RNAscope® VS Universal APAssay.

Deparaffinize FFPE sections

IMPORTANT! If you have not done so already, create a protocol for your instrument and print slide labels during this procedure. See pages 25–26.

1. Place slides in a Tissue-Tek® Slide Rack and submerge in the first xylene-containing clearing agent dish in the fume hood.
2. Incubate the slides in xylene for 5 MIN at RT. Agitate the slides by occasionally lifting the slide rack up and down in the clearing agent dish.
3. Remove the slide rack from the first xylene-containing dish and immediately place in the second xylene-containing clearing agent dish in the fume hood.
4. Repeat Step 2.
5. Remove the slide rack from the second xylene-containing dish and immediately place in the staining dish containing 100% EtOH.
6. Incubate the slides in 100% EtOH for 1 MIN at RT with agitation.
7. Repeat Step 6 with fresh 100% EtOH.
8. Remove the slides from the rack, and place on absorbent paper with the section face-up. Air dry for 5 MIN at RT.
9. While slides are drying, place printed labels on the slides.

IMPORTANT! Labels must be in place prior to the next section.

10. Insert the slides into a Tissue-Tek® Slide Rack and proceed to condition the slides.
Pretreat the slides

Begin heating 1X Target Retrieval Buffer while FFPE slides are baking at 60°C or during the previous section.

**IMPORTANT!** Do not boil 1X Target Retrieval more than 30 MIN before use.

1. Heat 1X Target Retrieval Buffer to 98–104°C:
   a. Place the beaker containing 1X Target Retrieval Buffer on the hot plate. Cover the beaker with foil and turn the hot plate on high for 10–15 MIN.
   b. Once 1X Target Retrieval Buffer reaches a slow boil (98–104°C), turn the hot plate to a lower setting to maintain the correct temperature. Check the temperature with a thermometer.

2. With a pair of forceps very slowly submerge the slide rack containing the slides into the boiling 1X Target Retrieval Buffer solution. Cover the beaker with foil and boil the slides for the amount of time specified in the following table:

<table>
<thead>
<tr>
<th>Tissue Type</th>
<th>Off Line Target Retrieval Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain and spinal cord</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Breast cancer</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Cell lines</td>
<td>10 MIN</td>
</tr>
<tr>
<td>Colon</td>
<td>15 MIN</td>
</tr>
<tr>
<td>GI tract</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Head and neck cancer</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Heart</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Kidney</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Liver</td>
<td>30 MIN</td>
</tr>
<tr>
<td>Lung</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Lymphoma</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Placenta</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Prostate</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Skin</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Stomach</td>
<td>15 MIN</td>
</tr>
<tr>
<td>Thymus</td>
<td>10 MIN</td>
</tr>
<tr>
<td>Tonsil</td>
<td>10 MIN</td>
</tr>
<tr>
<td>Xenograft derived from cell lines</td>
<td>7 MIN</td>
</tr>
<tr>
<td>Xenograft derived from primary tumor</td>
<td>15 MIN</td>
</tr>
</tbody>
</table>

3. Immediately transfer the hot slide rack from the 1X Target Retrieval Buffer to a staining dish containing distilled water. Do not let the slides cool in Target Retrieval.
4. Wash slides 3–5 times by moving the Tissue-Tek® Slide Rack up and down in the distilled water.
5. Repeat Step 4 with fresh distilled water.
6. Proceed directly to **Load the reagents** on page 29.
Run the RNAscope® VS Universal AP Assay

Materials required

- Prepared slides
- Prepared instrument reagents
- Prepared detergent
- Distilled water
- Prepared dehydrating materials
- Tissue-Tek® Vertical 24 Slide Rack
- EcoMount
- Cover Glass, 24 mm x 50 mm

Load the reagents

1. Remove the nozzle caps from the filled dispensers and place each cap on the post located on the back of the dispenser.
2. If needed, remove any air bubbles at the nozzle tips by pushing down on the nozzle until the liquid reaches the tip of the nozzle or forms a small meniscus at the tip of the nozzle
   **IMPORTANT!** Do not dispense any drops as this could compromise your drop inventory.
3. Load dispensers onto the reagent racks.
4. Remove the yellow locking ring from the dispensers in the prefilled mRNA RED Detection Kit. Refer to the instructions provided by Ventana™ Medical Systems.
5. Load the reagent racks onto the reagent carousel.

Start the run

1. Click the **Ready** button.

   ![State Chart](chart.png)

2. Eject slide drawers.
3. Load each slide onto a heater pad with the label facing upward and inward. Ensure that the slides sit securely on the pads.
   **IMPORTANT!** Prior to loading the slides, ensure heater pads are completely dry. Wipe off any solution using laboratory tissue paper.
5. Click the **Running** button. Semi-automated assay will finish in ~8 HRS.

   ![State Chart](chart.png)

**IMPORTANT!** Before leaving the instrument unattended, ensure all reagents and slides are successfully registered and the instrument is running.
Prepare detergent

1. Prepare 200 mL of diluted detergent by adding 1 to 2 drops detergent to 200 mL distilled water in a container with a cap.
2. Mix well by inverting the container 4–5 times.
3. Add diluted detergent to a Tissue-Tek® Staining Dish.

   Note: Store diluted detergent at RT.

Prepare dehydrating reagents

IMPORTANT! Do not use deparaffinization solutions for dehydration.

- In a fume hood, fill a clearing agent dish with ~200 mL fresh xylene.

   Note: Ensure all containers remain covered when not in use.

Complete the run

1. After the run is complete, place nozzle caps back on the dispensers.
2. Store reagent racks at 4°C until next use.

Wash the slides

1. Submerge a Tissue-Tek® Slide Rack into the Tissue-Tek® Staining Dish containing 200 mL diluted detergent.
2. Open the instrument slide drawer and unload slides.
3. Decant solution on the slides into the slide drawer, then immediately load slides into the Tissue-Tek® Slide Rack submerged in detergent.
4. Rinse oil off the slides by moving the slide rack up and down in the dish 10 times.
5. Replace the detergent with distilled water and rinse slides by moving the slide rack up and down a minimum of 10 times.
6. Repeat Step 5 three to five times.

Mount the samples

1. Remove the slide rack from the staining dish and dry slides in a 60°C dry oven for 30 MIN.

   IMPORTANT! The RED substrate is alcohol sensitive. Do not dehydrate the slides in alcohol.

2. Cool the slides for 5 MIN at RT.
3. Briefly dip one slide into into fresh pure xylene and immediately place 1–2 drops of EcoMount on the slide before the xylene dries.

   IMPORTANT! Use the EcoMount mounting medium only.

4. Carefully place 24 mm x 50 mm coverslip over the tissue section. Avoid trapping air bubbles.
5. Repeat steps 2 and 3 for each slide.
6. Air dry slides for at least 5 MIN.
7. Proceed to Chapter 5. Evaluate the Results.
**Recommended guidelines**

We highly recommend following the guidelines for Cell Conditioning (Target Retrieval) and Protease conditions for:

- Any new or previously untested FFPE tissue types.
- Samples prepared differently than the protocol in *Chapter 3. Prepare and Pretreat Samples* on page 11.

1. Stain six representative slides using the positive and negative control probes according to the following table:

<table>
<thead>
<tr>
<th>Slide No.</th>
<th>Probe</th>
<th>Target Retrieval</th>
<th>Protease</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Positive control</td>
<td>10 MIN</td>
<td>24 MIN</td>
</tr>
<tr>
<td>2</td>
<td>Negative control</td>
<td>10 MIN</td>
<td>24 MIN</td>
</tr>
<tr>
<td>3</td>
<td>Positive control</td>
<td>10 MIN</td>
<td>16 MIN</td>
</tr>
<tr>
<td>4</td>
<td>Negative control</td>
<td>10 MIN</td>
<td>16 MIN</td>
</tr>
<tr>
<td>5</td>
<td>Positive control</td>
<td>15 MIN</td>
<td>16 MIN</td>
</tr>
<tr>
<td>6</td>
<td>Negative control</td>
<td>15 MIN</td>
<td>16 MIN</td>
</tr>
</tbody>
</table>

2. Evaluate staining and tissue morphology as in *Chapter 5. Evaluate the Results*, and determine which pretreatment condition yielded the highest positive control signal and lowest negative control signal. Using PPIB, positive control signal should have a staining score of 3 or higher, and the negative control signal should be 0.

3. Use the optimized pretreatment conditions to run the assay with the target probe.

4. If none of the conditions are satisfactory, contact technical support at support.acd@bio-techne.com.
Appendix B. Safety

## Chemical safety

### WARNING!

**GENERAL CHEMICAL HANDLING.** To minimize hazards, ensure laboratory personnel read and practice the general safety guidelines for chemical usage, storage, and waste provided below, and consult the relevant SDS for specific precautions and instructions:

- Read and understand the Safety Data Sheets (SDSs) before you store, handle, or work with any chemicals or hazardous materials. To obtain SDSs, see https://acdbio.com/technical-support/user-manuals.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing).
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood).
- Characterize (by analysis if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure that the waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.
- **IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

## Biological hazard safety

### WARNING!

**BIOHAZARD.** Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Follow all applicable local, state/provincial, and/or national regulations. Wear appropriate protective equipment, which includes but is not limited to: protective eyewear, face shield, clothing/lab coat, and gloves. All work should be conducted in properly equipped facilities using the appropriate safety equipment (for example, physical containment devices). Individuals should be trained according to applicable regulatory and company/institution requirements before working with potentially infectious materials. Read and follow the applicable guidelines and/or regulatory requirements in the following:

### In the U.S.:

- U.S. Department of Health and Human Services guidelines published in Biosafety in Microbiological and Biomedical Laboratories found at: https://www.cdc.gov/biosafety/
- Your company’s/institution’s Biosafety Program protocols for working with/handling potentially infectious materials.
• Additional information about biohazard guidelines is available at:
  https://www.cdc.gov/biosafety/

In the EU:

• Check local guidelines and legislation on biohazard and biosafety precaution and refer to the best practices published in the World Health Organization (WHO) Laboratory Biosafety Manual, third edition, found at:

• Information about the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH) can be found at: https://echa.europa.eu/regulations/reach
Documentation and Support

Obtaining SDSs

Safety Data Sheets (SDSs) are available at: https://acdbio.com/technical-support/user-manuals. For the SDSs of chemicals not distributed by Advanced Cell Diagnostics, contact the chemical manufacturer.

Obtaining support

For the latest services and support information, go to: https://acdbio.com/technical-support/support-overview.

At the website, you can:

- Access telephone and fax numbers to contact Technical Support and Sales facilities.
- Search through frequently asked questions (FAQs).
- Submit a question directly to Technical Support.
- Search for user documents, SDSs, application notes, citations, training videos, and other product support documents.
- Find out information about customer training events.

Contact information

Advanced Cell Diagnostics, Inc.
7707 Gateway Blvd Suite 200
Newark, CA 94560
Toll Free: 1-877-576-3636
Direct: 1-510-576-8800
Fax: 1-510-576-8801
Information: info.acd@bio-techne.com
Orders: orders.acd@bio-techne.com
Support Email: support.acd@bio-techne.com

Limited product warranty

Advanced Cell Diagnostics, Inc. and/or its affiliate(s) warrant their products as set forth in the ACD General Terms and Conditions of Sale found on the ACD website. If you have any questions, please contact Advanced Cell Diagnostics at https://acdbio.com/about/contact.